## AMENDMENTS TO THE CLAIMS

1. (Currently Amended) A method for *in vivo* down-regulation of osteoprotegerin ligand (OPGL) activity in an animal, the method comprising effecting presentation to the animal's immune system of an immunogenically effective amount of an modified OPGL polypeptide analogue having general formula I:

$$(MOD_1)_{s1}(OPGL_{e1})_{n1}(MOD_2)_{s2}(OPGL_{e2})_{n2}...(MOD_x)_{sx}(OPGL_{ex})_{nx}$$
 (I)

-where  $OPGL_{e1}$ - $OPGL_{ex}$  are x B-cell epitope containing subsequences of OPGL which independently are identical or non-identical and which optionally contain foreign side groups, x is an integer  $\geq 3$ , n1-nx are x integers  $\geq 0$  of which at least one is  $\geq 1$ ,  $MOD_1$ - $MOD_x$  are x modifications introduced between the preserved B-cell epitopes, and  $s_1$ - $s_x$  are x integers  $\geq 0$  of which at least one is  $\geq 1$  if no optional side groups are introduced in the  $OPGL_e$  sequences, whereby the animal's own OPGL is down-regulated due to binding thereof to antibodies induced by immunization with the modified OPGL polypeptide analogue,

OPGL being a protein which acts as an osteoclast differentiation factor and which has an amino acid sequence as set forth in SEQ ID NO: 2 for human OPGL and in SEQ ID NOs: 4 and 6 for murine OPGL.

- 2. (Cancelled)
- 3. (Currently Amended) The method according to claim 1, wherein the analogue modified OPGL polypeptide comprises at least one foreign T helper lymphocyte epitope (T<sub>H</sub> epitope).
- 4. (Cancelled)

- 5. (Currently Amended) The method according to claim 1, wherein the <u>modified OPGL</u> polypeptide analogue includes an amino acid substitution in or deletion in or insertion in or addition to the OPGL polypeptide sequence, or any combination thereof.
- 6. (Cancelled)
- 7. (Cancelled)
- 8. (Currently Amended) The method according to claim 1, wherein the <u>modified OPGL</u> polypeptide <u>analogue-includes</u> a duplication of at least one OPGL B-cell epitope.
- 9. (Previously Presented) The method according to claim 3, wherein the foreign T-cell epitope is immunodominant in the animal.
- 10. (Previously Presented) The method according to claim 9, wherein the foreign T-cell epitope is capable of binding to a large proportion of MHC Class II molecules.
- 11. (Original) The method according to claim 10, wherein the at least one foreign T-cell epitope is selected from a natural T-cell epitope and an artificial MHC-II binding peptide sequence.
- 12. (Currently Amended) The method according to claim 11, wherein the natural T-cell epitope is selected from a Tetanus toxoid epitope. [such as P2 or P30 (SEQ ID NOs: 34 and 35, respectively)], a diphtheria toxoid epitope, an influenza virus hemagluttinin epitope, and a P. falciparum CS epitope.

Claims 13-16 (Cancelled)

17. (Currently Amended) The method according to claim 1, wherein the <u>modified OPGL</u> polypeptide <del>analogue contains a modification in any one of positions 171-193, any one of</del>

positions 199-219, any one of positions 222-247, any one of positions 257-262, or in any one of positions 286-317, the amino acid numbering conforming with that of SEQ ID NO: 2.

- 18. (Original) The method according to claim 17, wherein the modification comprises a substitution of at least one amino acid sequence within a position defined in claim 17 with an amino acid sequence of equal or different length which contains a foreign T<sub>H</sub> epitope.
- 19. (Previously Presented) The method according to claim 18, wherein the amino acid sequence containing the foreign T<sub>H</sub> epitope substitutes amino acids 257-262 and/or 289-303 and/or 222-243 in SEQ ID NO: 2 or in a polypeptide where a cysteine corresponding to Cys-221 of SEQ ID NO: 2 has been substituted with Ser.
- 20. (Currently Amended) The method according to\_claim 1, wherein presentation to the immune system is effected by having at least two copies of the <u>modified OPGL</u> polypeptide analogue-covalently or non-covalently linked to a carrier molecule capable of effecting presentation of multiple copies of antigenic determinants.
- 21. (Currently Amended) The method according to claim 1, wherein the <u>modified OPGL</u> polypeptide <del>analogue</del> has been formulated with an adjuvant which facilitates breaking of autotolerance to autoantigens.
- 22. (Currently Amended) The method according to\_claim 1, wherein an effective amount of the <u>modified OPGL</u> polypeptide <del>analogue</del> is administered to the animal via a route selected from the parenteral route selected from the group consisting of the intracutaneous, the subcutaneous, and the intramuscular routes; the peritoneal route; the oral route; the buccal route; the sublinqual route; the epidural route; the spinal route; the anal route; and the intracranial route.
- 23. (Currently Amended) The method according to claim 22, wherein the effective amount is between 0.5 µg and 2,000 µg of the modified OPGL polypeptide-analogue.

24. (Currently Amended) The method according to claim 22, wherein the <u>modified OPGL</u> polypeptide <del>analogue</del> is contained in a virtual lymph node (VLN) device.

Claims 25-27 (Cancelled)

28. (Currently Amended) The method according to claim 22, wherein the modulated OPGL polypeptide is administered or introduced which includes at least 1 one administration/introduction per year, such as at least 2, at least 3, at least 4, at least 6, orand at least 12-administrations/introductions times per year.

Claims 29-56 (Cancelled)

- 57. (Previously Presented) The method according to claim 1, wherein the animal is a human being.
- 58. (Previously Presented) The method according to claim 12, wherein the Tetanus toxoid epitope is a P2 or P30 epitope.